Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
1	774	544/58:2, 514/227.8	USPAT	OR	OFF	2005/08/30 16:05
L2	426	544/58.2, 514/227.8	US-PGPUB	OR	OFF	2005/08/30 16:05
L4	1200	544/58:2, 514/227:8	US-PGPUB;	OR	OFF	2005/08/30 16:05

10/766,122

Page 4

=> d l1 L1 HAS NO ANSWERS STR

Structure attributes must be viewed using STN Express query preparation.

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SAMPLE SEARCH INITIATED 12:45:39 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 0 TO ITERATE

100.0% PROCESSED

0 ITERATIONS

0 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS:

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FULL SEARCH INITIATED 12:45:46 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 10 TO ITERATE

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2 ANSWERS

SEARCH TIME: 00.00.01

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COST IN U.S. DOLLARS SINCE FILE TOTAL ENTRY SESSION

<08/30/2005> Habte 10/766,122

Page 5

FULL ESTIMATED COST

161.33 161.54

FILE 'CAPLUS' ENTERED AT 12:45:50 ON 30 AUG 2005 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS.

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This file contains CAS Registry Numbers for easy and accurate substance identification.

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L4 ANSWER 1 OF 5 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2005:23200 CAPLUS

DOCUMENT NUMBER: 112:13463

ITYENTOR (S): 12:13463

INVENTOR (S): Preparation of 2-phenyl-N-(pyridin-3-yl)-Nmethylisobutyramide derivatives as dual NKI/NK3
antagonists for treating schizophrenia
antagonists for treating schizophrenia
Schnider, Patricks Sleight, Andrews Stadler, Heinz
Schnider, Patricks Sleight, Andrews Stadler, Heinz
PATENT ASSIGNEE(S): F. Hoffmann-La Roche A.-G., Switz.
PCT Int. Appl., 374 pp.
CODEN: PIXXU2

DOCUMENT TYPE: Patent
LANGUAGE: Patent
English
FAMILY ACC. NUM. COUNT: 1

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

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PATENT NO.						KIND DATE				APPL	ICAT		DATE						
	WO	WO 2005002577			A1 20050			0113		WO 2	004-	29	20040625						
		₩:	AE,	λG,	λL,	AM,	AŤ,	AU,	λZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
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			GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	
			LK.	LR,	LS,	LT,	LU,	LV,	HΑ,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,	
			NO.	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	5K,	SL,	SY,	
			TJ,	TH,	TN,	TR.	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW	
		RW:	BW.	GH,	GM,	KE,	LS,	MW.	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	
			AZ.	BY,	KG,	KZ,	MD,	RU,	ŦJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	
			EE.	ES.	FI,	FR.	GB,	GR,	HU,	IR,	IT,	LU,	HC,	NL,	PL,	PT,	RO,	SE,	
			SI.	SX.	TR.	BF.	BJ,	CF,	CG,	CI,	CH,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	
			SN,	TD.	TG														
US 2005090533						A1 2005042			0428		US 2	004-	8847	07	20040702				
PRIORITY APPLN. INFO.:										EP 2	003-	1451	3		A 2	0030	703		
٥	THER SO	OURCE	(5):			MARPAT 142:134463													
G	I																		

ANSWER 1 OF 5 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

6

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 1 OF 5 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

$$\begin{array}{c|c} & & & \\ & & & \\ R^5 R^4 N & & & \\ \end{array} \begin{array}{c} & & \\ N & & \\ \end{array} \begin{array}{c} & \\ N & \\ \end{array} \begin{array}{c} & \\ N & \\ \end{array} \begin{array}{c} \\ N^3 & \\ \end{array} \begin{array}{c} \\ \end{array} \begin{array}{c} \\ R^3 & \\ \end{array} \begin{array}{c} \\ \end{array} \begin{array}{c} \\ \end{array} \begin{array}{c} \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \end{array} \begin{array}{c} \\ \\ \\ \\ \\$$

The invention is directed to the use of compds. of formula I [wherein Rl = (un) substituted aryl) R2, R3 = independently H, halo, alkyl, alkowy, OCHZP, OCHZP, OCF3, or CF3, R4, R5 = independently H, CHO, (CH2) OS(0)p-alkyl, etc. or 0-3p p 0-2 or R4NR5 form an (un) substituted ring with - (CH2) 3-5-, -(CH2)1, 2, 3-0-(CH2)2-, -CH2CH:CHCH2-, etc.)) and their pharmaceutically active acid addition salts as dual neurokinin NKI/NK3 antagonists useful in the treatment of schizophrania. The invention discloses 421 prepars. of title compds For example, II was prepared, in 2 steps, by acylation of N-(6-chloro-4-(2-chlorophenyl)pyridin-3-yl] methylamine (preparation given) with 2-(3,5-dichlorophenyl)-2-methylpropancyl chloride (preparation given) and amination with (1)-prolinol. II bound to NK1 and NK3 receptors with pKi value of 0.47 and 9.05, resp.
225641-09-6F, 2-[3,5-bis[trifluoromethyl)phenyl]-N-(4-(4-fluoro-2-methylphenyl)-6-(3-phydroxymethyl-1,1-dioxo-4-thiomorpholinyl)pyridin-3-yl]-N-methylisobutyramide
(Ri: PAC (Rharmacological activity); SPN (Synthetic preparation); USES (Uses)
(drug candidate; preparation of N-(pyridin-3-yl)-N-methylisobutyramide derivs. as dual NKI/KK3 antagonists for treating schizophrenia)
225641-09-6 CAPLUS
Benzenesetanide, N-[4-(4-fluoro-2-methylphenyl)-6-[3-(hydroxymethyl)-1,1-dioxido-4-thiomorpholinyl]-3-pyridinyl-N,s,a-trimethyl-3,5-bis(trifluoromethyl)- (SCI) (CA INDEX NAME)

00/ work

L4 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2005 ACS ON STN ACCESSION NUMBER: 2004:648391 CAPLUS DOCUMENT NUMBER: 141:174195

141:174195
A preparation of new crystalline modifications of 2-(3,5-bis-trifluoromethyl-phenyl)-N-[6-(1,1-dioxo-1A6-thiomorpholin-4-yl)-4(4-fluoro-2-methyl-phenyl)-pyridin-3-yl]-N-methylisobutyramide, useful as NKI receptor antagonists
HOffmann, Torsten: HOffmann-Emery, Fabienne; Nick, Soniar Schwitter, Urs; Waldmeier, Pius
F. Hoffmann-La Roche AG, Switz.
PCT Int. Appl., 27 pp.
CODEN: PIXEU2
Patent
English

INVENTOR (S):

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: English

LANGUAGE: FAMILY ACC, NUM. COUNT: PATENT INFORMATION:

PATENT NO.				KIND DATE				APPL	ICAT	DATE								
	wo	2004	0670	07		Al	-	2004	0812	1	WO 2	004-	EP 54	7		2	0040	123
		W:	AE,	AE,	AG,	AL,	AL,	AM,	AM,	AM.	AT.	AT.	AU.	AZ.	AZ.	BA.	BB.	BG.
								BY,										
			CU,	CU,	cz,	cz,	DE,	DE,	DK,	DK,	DM,	DZ,	EC.	EC,	EE.	EE.	EG.	ES,
								GE,										
			IS,	JP,	JP,	KE,	KE,	KG.	KG,	KP.	KP.	KP,	KR,	KR.	KZ.	KZ,	KZ.	LC.
			LK,	LR,	LS,	LS,	LT.	LU,	LV.	MA,	MD,	MD.	MG.	MK.	MN.	MW.	MX.	MX.
				MZ.														
	US	2004	1861	00		A1		2004	0923	1	US 2	004-	7661	22 _		2	0040	127
PRIO	RIT	APF	LN.	INFO	.:						EP Z	003-	2134		٠.	A 2	0030	131

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB The invention relates to a preparation of new crystalline modifications of 2-(3,5-bis-trifluoronethyl-phenyl)-N-[6-(1,1-dioxo-1A6-thiomorpholin-4-yl)-4(4-fluoro-2-methyl-phenyl)-pyridin-3-yl]-N-methyl-isobutyramide (I) characterized by X-ray diffraction and useful as NKI receptor antagonists. I was prepared via amidation of the obtained propionic scid derivative II by thiomorpholinylpyridine derivative III and subsequent 5-oxidation Four modifications of I were identified: 3

Crystalline (A, B, C) and one amorphous. Form A demonstrated the highest bioavailability among the three crystalline polymorphs A, B, and C.

1474026-04-5P

RL: PAC (Phermacological activity): PKT (Pharmacokinetics): PRP (Properties): SFN (Synthetic preparation): TRU (Therapeutic use): BIOL (Biological study): PREP (Preparation): USES (Uses) (preparation of new crystalline modifications of thiomorpholinylpyridine derivative,

useful as NKI receptor antagonists)

RN 474026-04-5 CAPIUS

C Benzenacctamide, N-[6-(1,1-dioxido-4-thiomorpholinyl)-4-{4-fluoro-2-methylphenyl)-3-pyridinyl]-N, a, a-trimethyl-3, 5-bis(trifluoromethyl)- (9CI) (CA INDEX NAME)

10/766,122

Page 7

ANSWER 2 OF 5 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)

ANSWER 3 OF 5 CAPLUS COPYRIGHT 2005 ACS on STW (Continued) detecting a polymorphism in the NXNA gene, to a pharmaceutical pack comprising neurokinin-1 (NX-1) receptor antagonists and instructions for administration of the drug to human beings tested for the polymorphisms as well as to a computer readable medium with the stored sequence information for the polymorphisms in the NXNA gene.
474026-04-5

ATO228-04-5

RL: ANT (Analyte), PAC (Pharmacological activity), THU (Therapeutic use);

ANST (Analytical study), BIOL (Biological study), USES (Uses)

(NK-1 receptor antagonist, method for correlating preprotachykinin gene

(NKNA) polymorphisms with efficacy and compatibility of

pharmaceutically active compds., such as NK-1 receptor antagonists)

47025-04-5 CAPLUS

Benzeneacetamide, N-[6-(1,1-dioxido-4-thiomorpholinyl)-4-(4-fluoro-2-methylphenyl)-3-pyridinyl]-M.y.,a-trimethyl-3-5
bis(trifluoromethyl)- (9CI) (CA INDEX NAME)

L4 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:
DOCUMENT NUMBER:
139:160766
A method for correlating the preprotachykinin gene
(NUMA) polymorphisms with the efficacy and
compactibility of a pharmaceutically active compounds,
such as NK-1 receptor antagonists
Foernzier, Dorothee Hashimoto, Lara; Li, Jia; Luedin,
Eric; Sleight, Andrew; Vankan, Pierre
F. Hoffmann-La Roche A.-G., Switz.

COUNS! FIXON2

DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
FAMILY ACC. NUM. COUNT:
1 DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2003064685 A2 20030807 WO 2003-RFG30 20030123

WO 2003064685 A3 20031224

W. AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, BE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, HD, MG, MK, MM, MM, MK, MZ, CM, NZ, CM, PH, PL, PT, RO, RU, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VN, VU, ZA, ZM, ZV

RW: GH, GM, KE, LS, HW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TH, ATD, ES, SI, SK, TR, BF, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GM, GQ, GW, ML, MR, NE, SN, TD, TG

CA 2473128 AA 20030807 CA 2003-2473128 20030123

EP 1472377 A2 20041103 EP 2003-734685 20030123

ER: AT, BE, CH, DE, DIK, MS, FR, GB, GR, IT, LI, LU, HL, SE, MC, PT, IS, SI, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK

BR 2003007257 A 20041214 BR 2003-7257 20030123

JF 2005515788 T2 20050602 JF 2003-24575 20030123

PRIORITY APPLN. INFO:

EP 2002-1937 A 20020131

AB The present invention relates to a method for correlating single nucleotide polymorphisms in the preprotachykinin (NNMA) gene with the efficacy and compatibility of a pharmaceutically active compound administered to a human being. The invention further relates to a method for determining the efficacy and compatibility of a pharmaceutically active compound administered to a human being which method comprises determining at least on determining specific single nucleotide polymorphisms in the NNNA gene. Said methods are based on determining specific single nucleotide polymorphism in the NNNA gene. Said methods are based on determining specific single nucleotide polymorphisms in the NNNA gene. Said methods are based on determining specific single nucleotide polymorphisms in the NNNA gene. Said methods are based on determining specific single nucleotide polymorphisms in the NNNA gene. Said methods are APPLICATION NO. PATENT NO. KIND DATE DATE one single nucleotide polymorphism in the NXNA gene. Said methods are based on determining specific single nucleotide polymorphisms in the NXNA and determining the efficacy and compatibility of a pharmaceutically active compound in the human by reference to polymorphism in NKNA. The invention further relates to isolated nucleic acids comprising within their sequence the polymorphisms as defined herein, to nucleic acid primers and oligonucleotide probes capable of hybridizing to such nucleic acids and to a diagnostic kit comprising one or more of such primers and probes for

L4 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2005 ACS on STN
ACCESSION NUMBER: 2003:117823 CAPLUS
DOCUMENT NUMBER: 138:170243
TITLE: Preparation of 2-(3.5-bis-tri 138:170243

Preparation of 2-(3,5-bis-trifluoromethyl-phenyl)-N-[6-(1,1-dioxo-1λ6-thiomorpholin-4-yl)-4-(2-methyl or 4-fluoro-2-methyl substituted)phenyl-pyridin-3-yl]-N-methyl-isobutyramide as selective NKI antagonists Ballard, Theresa Maria; Hoffmann, Torsten; Poli, Sonia Maria; Schnider, Patrick; Sleight, Andrew F. Hoffmann-La Roche AG, Switz.

CODEN: PIXXD2

PRESENT INVENTOR(S): PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: Patent English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE A2 A3 WO 2003011860 WO 2003011860 20030213 WO 2002-EP8311 20020726 20030904

OTHER SOURCE(S): MARPAT 138:170243

The title compds. I [Rl = H. F] which may be used for the treatment of migraine, rheumatoid arthritis, asthma, bronchial byperreactivity, inflammatory bowel disease or for the treatment of disorders including Parkinson's disease, anxiety, depression, pain, headache, Altheimer's disease, multiple sclerosis, deman, altergic rhintis, Crohn's disease, ocular injury, ocular inflammatory diseases, psychosis, motion sickness, induced vomiting, emesis, urinary incontinence, psychoimmunol. or psychosomatic disorders, cancer, withdrawal symptoms of addictive drugs from opiates or nicotine, traumatic brain injury or benign prostatic hyperplasis, were prepared and formulated. E.g., a 8-step synthesis of I[Rl = H] (starting with 2-chloro-5-nitropyridine and thiomorpholine) which showed pKi of 8.9 for the human NKI receptor, was given.

476026-04-59
RL: PAC (Pharmacological activity) SFN (Synthetic preparation), THU (Therapautic use), BIOL (Biological study), PREP (Preparation) USES (Uses) AB

(Uses)
((Uses)
((Uses)
((Uses))
((Uses)

L4 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2005 ACS ON STN ACCESSION NUMBER: 2002:832668 CAPLUS DOCUMENT NUMBER: 137:337901 TITLE: Feparation and use of amide:

137:33/901
Preparation and use of amides as NK-1 receptor antagonists against benign prostatic hyperplasia Buser, Susanne; Ford, Anthony P. D. W.; Hoffmann, Torsten; Lenz, Barbara; Sleight, Andrew John; Vankan, INVENTOR(S):

Torsten, Lenz, Barbara; Sleight, Ar Pierre F. Hoffmann-La Roche A.-G., Switz. PCT Int. Appl., 45 pp. CODEN: PIXXD2 Patent PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE:

LANGUAGE: English

FAMILY ACC. NUM. COUNT:

ATEN	IT I	NFOR	MATI	ON:														
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	WO	2002	0854	58		A2		2002	1031	,	WO	2002-	EP 10	85		2	0020	202
	WO	2002	0854	58		A3		2003	1030									
		W:	AE,	AG,	AL,	AM,	AT,	AU,	AΖ,	BA,	BB	, BG,	BR,	BY,	BZ,	CA,	CH,	CN,
			co.	CR.	CU.	CZ.	DE.	DK.	DM.	DZ.	EC	, EE,	ES,	FI.	GB,	GD,	GE,	GH,
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	BR	2002	0091	51		Α		2004	0713		BR	2002- 2002- 2002-	9151			2	0020	202
	JP	2004	5299	31		T2		2004	0930		JP	2002-	5830	31		2	0020	202
	US	2003	0041	57		A1		2003	0102		US	2002-	7157	0		2	0020	208
	ZA	2003	0081	10		A		2005	0117		ZA	2003-	8110			2	0031	017
RIO	IT	APP	LN.	INFO	.:							2001-						
											WO	2002-	BP10	85		W 2	0020	202
THE	3 50	URCE	(S):			MAR	PAT	137:	3379	01								

ANSWER 5 OF 5 CAPLUS COPYRIGHT 2005 ACS on STN (Continued)
Use of an NK-1 receptor antagonist for the treatment or prevention of benign prostatic hyperplasia (BFH) is claimed. The preferred NK-1 receptor antagonists are compds, of the general formula [1, R = H, alkyl, alkoxy, halo, CF3; Rl = H, halo; RR1 = CH:CHCH:CH: NZ, RZ1 = H, halo; RC73, alkyl, alkoxy, cyanon RZR21 = CH:CHCH:CH: optionally substituted by 1-2 alkyl, halo, alkoxy, R3 = H, alkyl; R3R3C = cycloalkyl; R4 = H, N(RS)2, NRS(CH2) nA, cyclic tertiary amine, etc.; X = CONRS, (CH2) pO, NRS(CH2) pD, etc.; R5 = H, cycloalkyl; Ph, PhCH2, alkyl; n = 0-4; p = 1-3]. Preferred compds. are 2-(3,5-bis-trifluoromethyl-phenyl)-M-methyl-N-[6-(1,1-dioxo-1Ac-thiomorpholin-4-yl-4-o-tolyl-pyridin-3-yl)isobutyramide, 3-(3,5-bis-trifluoromethyl-phenyl)-M-methyl

n)
oxone were stirred 2 days at room temperature to give 2-(3,5-bistrifluoromethylphenyl)-N-[6-(1,1-dioxo-126-thiomorpholin-4-yl)-4-otolylpyridin-3-yl)-N-methylisobutyramide. 2-(3,5Bistrifluoromethylphenyl)-N-methyl-N-methyl-N-(6-morpholin-4-yl-4-otolylpyridin-3-yl)isobutyramide at 60 mg/kg/day orally in dogs reduced
prostate weight by 58% after 39 wk.
474026-04-59

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(preparation and use of amides as NK-1 receptor antagonists against benian

gn prostatic hyperplasia)
474026-04-5 CAPLUS
Banzaneacetamide, N-[6-(1,1-dioxido-4-thiomorpholiny1)-4-(4-fluoro-2-methylpheny1)-3-pyridiny1)-N,α,α-trimethyl-3,5-bis(trifluoromethyl)- (9CI) (CA INDEX NAME)